

# TRAITE DE COOPERATION EN MATIERE DE BREVETS

## PCT

### RAPPORT D'EXAMEN PRELIMINAIRE INTERNATIONAL

(article 36 et règle 70 du PCT)

Référence du dossier du déposant ou du mandataire <b>BIF 22127 PCT</b>	<b>POUR SUITE A DONNER</b> voir la notification de transmission du rapport d'examen préliminaire international (formulaire PCT/IPEA/416)	
Demande Internationale n° <b>PCT/FR00/00118</b>	Date du dépôt international (jour/mois/année) <b>20/01/2000</b>	Date de priorité (jour/mois/année) <b>21/01/1999</b>
Classification internationale des brevets (CIB) ou à la fois classification nationale et CIB <b>B01F17/00</b>		
Déposant <b>ARCANE INDUSTRIES et al.</b>		
<p>1. Le présent rapport d'examen préliminaire international, établi par l'administration chargée de l'examen préliminaire international, est transmis au déposant conformément à l'article 36.</p> <p>2. Ce RAPPORT comprend 5 feuilles, y compris la présente feuille de couverture.</p> <p><input type="checkbox"/> Il est accompagné d'ANNEXES, c'est-à-dire de feuilles de la description, des revendications ou des dessins qui ont été modifiées et qui servent de base au présent rapport ou de feuilles contenant des rectifications faites auprès de l'administration chargée de l'examen préliminaire international (voir la règle 70.16 et l'instruction 607 des Instructions administratives du PCT).</p> <p>Ces annexes comprennent feuilles.</p>		
<p>3. Le présent rapport contient des indications relatives aux points suivants:</p> <ul style="list-style-type: none"> <li>I <input checked="" type="checkbox"/> Base du rapport</li> <li>II <input type="checkbox"/> Priorité</li> <li>III <input type="checkbox"/> Absence de formulation d'opinion quant à la nouveauté, l'activité inventive et la possibilité d'application industrielle</li> <li>IV <input type="checkbox"/> Absence d'unité de l'invention</li> <li>V <input checked="" type="checkbox"/> Déclaration motivée selon l'article 35(2) quant à la nouveauté, l'activité inventive et la possibilité d'application industrielle; citations et explications à l'appui de cette déclaration</li> <li>VI <input type="checkbox"/> Certains documents cités</li> <li>VII <input checked="" type="checkbox"/> Irrégularités dans la demande internationale</li> <li>VIII <input checked="" type="checkbox"/> Observations relatives à la demande internationale</li> </ul>		

Date de présentation de la demande d'examen préliminaire internationale <b>18/08/2000</b>	Date d'achèvement du présent rapport <b>26.04.2001</b>
Nom et adresse postale de l'administration chargée de l'examen préliminaire international:  Office européen des brevets D-80298 Munich Tél. +49 89 2399 - 0 Tx: 523656 epmu d Fax: +49 89 2399 - 4465	Fonctionnaire autorisé  Haderlein, A N° de téléphone +49 89 2399 2095



# RAPPORT D'EXAMEN PRÉLIMINAIRE INTERNATIONAL

Demande internationale n° PCT/FR00/00118

## I. Base du rapport

1. En ce qui concerne les éléments de la demande internationale (*les feuilles de remplacement qui ont été remises à l'office récepteur en réponse à une invitation faite conformément à l'article 14 sont considérées dans le présent rapport comme "initialement déposées" et ne sont pas jointes en annexe au rapport puisqu'elles ne contiennent pas de modifications (règles 70.16 et 70.17)*):

**Description, pages:**

1-14 version initiale

### **Revendications. N°:**

1-10 version initiale

2. En ce qui concerne la **langue**, tous les éléments indiqués ci-dessus étaient à la disposition de l'administration ou lui ont été remis dans la langue dans laquelle la demande internationale a été déposée, sauf indication contraire donnée sous ce point.

Ces éléments étaient à la disposition de l'administration ou lui ont été remis dans la langue suivante: qui est:

- la langue d'une traduction remise aux fins de la recherche internationale (selon la règle 23.1(b)).
- la langue de publication de la demande internationale (selon la règle 48.3(b)).
- la langue de la traduction remise aux fins de l'examen préliminaire internationale (selon la règle 55.2 ou 55.3).

3. En ce qui concerne les **séquences de nucléotides ou d'acide aminés** divulguées dans la demande internationale (le cas échéant), l'examen préliminaire internationale a été effectué sur la base du listage des séquences :

- contenu dans la demande internationale, sous forme écrite.
- déposé avec la demande internationale, sous forme déchiffrable par ordinateur.
- remis ultérieurement à l'administration, sous forme écrite.
- remis ultérieurement à l'administration, sous forme déchiffrable par ordinateur.
- La déclaration, selon laquelle le listage des séquences par écrit et fourni ultérieurement ne va pas au-delà de la divulgation faite dans la demande telle que déposée, a été fournie.
- La déclaration, selon laquelle les informations enregistrées sous déchiffrable par ordinateur sont identiques à celles du listages des séquences Présenté par écrit, a été fournie.

#### 4. Les modifications ont entraîné l'annulation :

- de la description, pages :
- des revendications, n°s :
- des dessins, feuilles :

**RAPPORT D'EXAMEN  
PRÉLIMINAIRE INTERNATIONAL**

Demande internationale n° PCT/FR00/00118

5.  Le présent rapport a été formulé abstraction faite (de certaines) des modifications, qui ont été considérées comme allant au-delà de l'exposé de l'invention tel qu'il a été déposé, comme il est indiqué ci-après (règle 70.2(c)) :

*(Toute feuille de remplacement comportant des modifications de cette nature doit être indiquée au point 1 et annexée au présent rapport)*

6. Observations complémentaires, le cas échéant :

**V. Déclaration motivée selon l'article 35(2) quant à la nouveauté, l'activité inventive et la possibilité d'application industrielle; citations et explications à l'appui de cette déclaration**

1. Déclaration

Nouveauté	Oui : Revendications 3,6 Non : Revendications 1,2,4,5,7-10
Activité inventive	Oui : Revendications 3,6 Non : Revendications
Possibilité d'application industrielle	Oui : Revendications 1-10 Non : Revendications

2. Citations et explications  
**voir feuille séparée**

**VII. Irrégularités dans la demande internationale**

Les irrégularités suivantes, concernant la forme ou le contenu de la demande internationale, ont été constatées :  
**voir feuille séparée**

**VIII. Observations relatives à la demande internationale**

Les observations suivantes sont faites au sujet de la clarté des revendications, de la description et des dessins et de la question de savoir si les revendications se fondent entièrement sur la description :  
**voir feuille séparée**

**RAPPORT D'EXAMEN****PRELIMINAIRE INTERNATIONAL - FEUILLE SEPARÉE**

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**Concernant les points V et VIII****1. Clarté**

La demande ne remplit pas la condition visée à l'article 6 PCT les revendications n'étant pas claires. Les raisons sont les suivantes.

- 1.1 L'expression "essentiellement" (revendication 1) est vague (cf. Directives III-4.5). Il serait préférable d'utiliser une expression concrète telle que "au moins 90% en poids" (voir p.2, dernier paragraphe de la demande).
- 1.2 La présence d'un agent actif hydrophobe dans la composition semble indispensable (cf. p. 2, 2ième alinéa ; p.8, I.17-19). Or la revendication 1 comprend aussi des mélanges ne contenant que de l'éther de glycol et des traces d'une N-alkyl pyrrolidone (cf. "au moins 10% en poids d'un agent actif ... ou moins de 10% d'une N-alkyl pyrrolidone"). Par conséquent, la revendication 1 n'est pas claire et ne répond pas à l'exigence de l'article 6 PCT (cf. les Directives III-4.3 : **caractéristique indispensable manquante**).
- 1.3 L'expression "diluable" (revendication 1) est vague parce qu'il n'est pas clair dans quel solvant la composition serait diluable. Dans la revendication 8 il est fait également référence à la dilution tandis qu'il n'est pas clair au moyen de quel solvant elle se fait.

**2. Nouveauté**

- 2.1 Le document **D1 : WO 91 00893 A** (cité dans la demande) décrit à sa page 4 une composition comprenant [essentiellement] :
  - 0% à 60 % en poids d'un dérivé terpénique (agent actif hydrophobe) et
  - 5% à 60% en poids d'éthers de glycol.

L'objet de la revendication 1 n'est donc pas nouveau (art. 33(2) PCT).

- 2.2 Les dérivés terpéniques possèdent un indice KB supérieur à 30 et notamment à 40 (cf. p.3,I.27-31 de la demande). Par conséquent, l'objet de la revendication 2 n'est pas nouveau (article 33(2) PCT).

**RAPPORT D'EXAMEN**

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**PRELIMINAIRE INTERNATIONAL - FEUILLE SEPARÉE**

2.3 Les éthers utilisés dans le document D1 sont notamment :

- l'éther méthylique du dipropylène glycol, l'éther n-propyle du propylène, l'éther n-propyle du dipropylène glycol et l'éther n-butylique du propylène glycol.

Par conséquent, l'objet de la revendication 5 n'est pas nouveau (art. 33(2) PCT).

2.4 L'objet des revendications 7 et 8 ne semble pas contenir une caractéristique qui pourrait remédier à l'objection de la nouveauté manquante (art. 33(2) PCT).

2.5 En autant que les revendications puissent être comprises (cf. l'alinéa "clarté" plus haut), les objets des revendications indépendantes de procédé et d'utilisation 9 et 10 respectivement ne sont pas nouveaux en vue du document D1 (art. 33(2) PCT).

3. Activité inventive

En autant que les revendications puissent être comprises (cf. l'alinéa "clarté" plus haut), le problème technique qui est à la base des revendications 3 et 6 est de réduire la quantité de N-alkyl pyrrolidone présente dans l'émulsion. Une émulsion comprenant au moins 10 % en poids d'un agent actif et deux éthers de glycol choisis selon le dernier alinéa de la revendication 1 est absente dans l'état de la technique disponible. Ceci étant donné, il n'est pas possible de prétendre qu'une modification de l'émulsion connu par le document D1 conduisant à l'objet des revendications 3 ou 6 est suggérée au vu de cet état de la technique disponible (art. 33(3) PCT).

Concernant le point VII

4. Contrairement à la règle 6.3 PCT, certaines des caractéristiques connues en combinaison dans l'état de la technique le plus proche (D1) figurent dans la partie caractérisante de la revendication indépendante 1.

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PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference BIF 22127 PCT	FOR FURTHER ACTION	See Notification of Transmittal of International Preliminary Examination Report (Form PCT/IPEA/416)
International application No. PCT/FR00/00118	International filing date (day/month/year) 20 January 2000 (20.01.00)	Priority date (day/month/year) 21 January 1999 (21.01.99)
International Patent Classification (IPC) or national classification and IPC B01F 17/00		
Applicant	ARCANE INDUSTRIES	

1. This international preliminary examination report has been prepared by this International Preliminary Examining Authority and is transmitted to the applicant according to Article 36.
2. This REPORT consists of a total of <u>5</u> sheets, including this cover sheet.
<input type="checkbox"/> This report is also accompanied by ANNEXES, i.e., sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT). These annexes consist of a total of _____ sheets.
3. This report contains indications relating to the following items:
I <input checked="" type="checkbox"/> Basis of the report
II <input type="checkbox"/> Priority
III <input type="checkbox"/> Non-establishment of opinion with regard to novelty, inventive step and industrial applicability
IV <input type="checkbox"/> Lack of unity of invention
V <input checked="" type="checkbox"/> Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement
VI <input type="checkbox"/> Certain documents cited
VII <input type="checkbox"/> Certain defects in the international application
VIII <input type="checkbox"/> Certain observations on the international application

Date of submission of the demand 18 August 2000 (18.08.00)	Date of completion of this report 26 April 2001 (26.04.2001)
Name and mailing address of the IPEA/EP	Authorized officer
Facsimile No.	Telephone No.

## INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/FR00/00118

## I. Basis of the report

1. With regard to the **elements** of the international application:\* the international application as originally filed the description:

pages \_\_\_\_\_, 1-14, as originally filed

pages \_\_\_\_\_, filed with the demand

pages \_\_\_\_\_, filed with the letter of \_\_\_\_\_

 the claims:

pages \_\_\_\_\_, 1-10, as originally filed

pages \_\_\_\_\_, as amended (together with any statement under Article 19

pages \_\_\_\_\_, filed with the demand

pages \_\_\_\_\_, filed with the letter of \_\_\_\_\_

 the drawings:

pages \_\_\_\_\_, as originally filed

pages \_\_\_\_\_, filed with the demand

pages \_\_\_\_\_, filed with the letter of \_\_\_\_\_

 the sequence listing part of the description:

pages \_\_\_\_\_, as originally filed

pages \_\_\_\_\_, filed with the demand

pages \_\_\_\_\_, filed with the letter of \_\_\_\_\_

2. With regard to the **language**, all the elements marked above were available or furnished to this Authority in the language in which the international application was filed, unless otherwise indicated under this item.

These elements were available or furnished to this Authority in the following language \_\_\_\_\_ which is:

 the language of a translation furnished for the purposes of international search (under Rule 23.1(b)). the language of publication of the international application (under Rule 48.3(b)). the language of the translation furnished for the purposes of international preliminary examination (under Rule 55.2 and/or 55.3).3. With regard to any **nucleotide and/or amino acid sequence** disclosed in the international application, the international preliminary examination was carried out on the basis of the sequence listing: contained in the international application in written form. filed together with the international application in computer readable form. furnished subsequently to this Authority in written form. furnished subsequently to this Authority in computer readable form. The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished. The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished.4.  The amendments have resulted in the cancellation of: the description, pages \_\_\_\_\_ the claims, Nos. \_\_\_\_\_ the drawings, sheets/fig \_\_\_\_\_5.  This report has been established as if (some of) the amendments had not been made, since they have been considered to go beyond the disclosure as filed, as indicated in the Supplemental Box (Rule 70.2(c)).\*\*

\* Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rule 70.16 and 70.17).

\*\* Any replacement sheet containing such amendments must be referred to under item 1 and annexed to this report.

**INTERNATIONAL PRELIMINARY EXAMINATION REPORT**International application No.  
PCT/FR 00/00118**V. Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement****1. Statement**

Novelty (N)	Claims	3, 6	YES
	Claims	1, 2, 4, 5, 7-10	NO
Inventive step (IS)	Claims	3, 6	YES
	Claims		NO
Industrial applicability (IA)	Claims	1-10	YES
	Claims		NO

**2. Citations and explanations**

See separate sheet.

**INTERNATIONAL PRELIMINARY EXAMINATION REPORT**

International application No.  
PCT/FR 00/00118

**VIII. Certain observations on the international application**

The following observations on the clarity of the claims, description, and drawings or on the question whether the claims are fully supported by the description, are made:

See separate sheet.

**Supplemental Box**

(To be used when the space in any of the preceding boxes is not sufficient)

Continuation of: separate sheet

**Concerning Boxes V and VIII**1. Clarity

The application does not fulfil the requirements of PCT Article 6, as the claims are unclear for the following reasons:

- 1.1 The term "mainly" (Claim 1) is vague (cf. the PCT Guidelines III-4.5). It would be preferable to use a specific expression such as "at least 90% by weight" (see the application, page 2, the last paragraph).
- 1.2 The presence of a hydrophobic active agent in the composition appears to be essential (cf. page 2, the second paragraph; page 8, lines 17-19). However, Claim 1 also includes mixtures that contain only glycol ether and traces of an N-alkyl pyrrolidone (cf. "at least 10% by weight of a hydrophobic active agent ... or less than 10% of an N-alkyl pyrrolidone"). As a result, Claim 1 is unclear and does not fulfil the requirements of PCT Article 6 (cf. the PCT Guidelines III-4.3: **missing essential feature**).
- 1.3 The term "dilutable" (Claim 1) is vague because it is unclear in which solvent the composition is allegedly dilutable. Reference is also made to dilution in Claim 8, yet it is unclear which solvent is used to perform said dilution.

**Supplemental Box**

(To be used when the space in any of the preceding boxes is not sufficient)

Continuation of: separate sheet

2. Novelty

2.1 Document **WO 91 00893 A (D1)** (cited in the application) describes a composition (page 4) containing [mainly]:

- 0% to 60% by weight of a terpene derivative (hydrophobic active agent) and
- 5% to 60% by weight of glycol ethers.

It follows that the subject matter of Claim 1 is not novel (PCT Article 33(2)).

2.2 The terpene derivatives have a KB index of more than 30 and, in particular, more than 40 (cf. page 3, lines 27-31 of the application). As a result, the subject matter of Claim 2 is not novel (PCT Article 33(2)).

2.3 The ethers used in document D1 are, in particular:

- dipropylene glycol methyl ether, propylene n-propyl ether, dipropylene glycol n-propyl ether and propylene glycol n-butyl ether.

It follows that the subject matter of Claim 5 is not novel (PCT Article 33(2)).

**INTERNATIONAL PRELIMINARY EXAMINATION REPORT**International application No.  
PCT/FR 00/00118**Supplemental Box**

(To be used when the space in any of the preceding boxes is not sufficient)

Continuation on: separate sheet

2.4 The subject matter of Claims 7 and 8 does not appear to contain any features that could obviate the objection to a lack of novelty (PCT Article 33(2)).

2.5 In so far as the claims can be understood (cf. the paragraph entitled "clarity" above), the subject matter of independent method Claim 9 and use Claim 10 is not novel over document D1 (PCT Article 33(2)).

3. Inventive step

In so far as the claims can be understood (cf. the paragraph entitled "clarity" above), the technical problem on which Claims 3 and 6 are based is that of reducing the amount of N-alkyl pyrrolidone in the emulsion. An emulsion containing at least 10% by weight of an active agent and two glycol ethers selected according to the last paragraph of Claim 1 is not included in the available prior art. In view of this, it is impossible to assert that said available prior art suggests a modification of the emulsion known from D1 that leads to the subject matter of Claims 3 or 6 (PCT Article 33(3)).

**INTERNATIONAL PRELIMINARY EXAMINATION REPORT**

International application No.  
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**VII. Certain defects in the international application**

The following defects in the form or contents of the international application have been noted:

Contrary to the requirements of PCT Rule 6.3, some of the features known in combination from the closest prior art (D1) are included in the characterising portion of independent Claim 1.



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5 :		A2	(11) International Publication Number:	WO 91/00893
C09D			(43) International Publication Date:	24 January 1991 (24.01.91)

(21) International Application Number: PCT/US90/03757

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## (30) Priority data:

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475,364	5 February 1990 (05.02.90)	US

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(81) Designated States: AT, AT (European patent), AU, BB, BE (European patent), BF (OAPI patent), BG, BJ (OAPI patent), BR, CA, CF (OAPI patent), CG (OAPI patent), CH, CH (European patent), CM (OAPI patent), DE\*, DE (European patent)\*, DK, DK (European patent), ES, ES (European patent), FI, FR (European patent), GA (OAPI patent), GB, GB (European patent), HU, IT (European patent), JP, KP, KR, LK, LU, LU (European patent), MC, MG, ML (OAPI patent), MR (OAPI patent), MW, NL, NL (European patent), NO, RO, SD, SE, SE (European patent), SN (OAPI patent), SU, TD (OAPI patent), TG (OAPI patent).

## Published

*Without international search report and to be republished upon receipt of that report.*

AA

(54) Title: PAINT STRIPPER AND VARNISH REMOVER COMPOSITIONS AND RELATED METHODS

## (57) Abstract

The present invention relates to non-toxic, non-flammable, safe compositions employing primarily biodegradable components for use in removing and stripping paint, varnishes and stains from hard and/or flexible surfaces. The compositions generally avoid the use of methylene chloride, methanol, toluol or acetone. Paint stripper compositions of the present invention have flash points above about 120 °F and an absence of low-boiling VOC's comprising: (a) about 0 % to about 60 % by weight of a terpene compound containing at least 10 carbon atoms; (b) about 10 % to about 69 % by weight of N-methyl pyrrolidone or its equivalent; and (c) about 0 % to about 20 % by weight of an emulsifying surfactant. Compositions of the present invention may further comprise one or more of the following: about 5 % to about 15 % by weight of a solvent extender or bulking agent, about 5 % to about 60 % of a high flash point, high K<sub>B</sub> (kauri-butanol) solvent selected from the group consisting of ethylene carbonate, propylene carbonate, butylene carbonate, glycols and glycol ethers, about 0 % to about 45 % water, from 0 % to about 8% by weight of a thixotropic or thickening agent, preferably an organoclay thickening agent, for example, steralkonium hectorite, about 2 % to about 20 % by weight of an odor masking component, for example d'limonene and mixtures, thereof. In the organoclay containing compositions according to the present invention, about 10 % to about 45 %, preferably about 30 to 45 % of a polar solvent, especially including water, ethanol, and mixtures of water and ethanol. Methods of stripping coating from surfaces and methods of making these compositions are also described.

## DESIGNATIONS OF "DE"

Until further notice, any designation of "DE" in any international application whose international filing date is prior to October 3, 1990, shall have effect in the territory of the Federal Republic of Germany with the exception of the territory of the former German Democratic Republic.

### FOR THE PURPOSES OF INFORMATION ONLY

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				US	United States of America

## Paint Stripper and Varnish Remover Compositions and Related Methods

### Field of the Invention

The present invention relates to non-toxic, non-flammable, safe compositions for use in removing and stripping paint, varnishes and stains from hard and/or flexible surfaces. In most instances, the compositions avoid the use of methylene chloride, methanol, toluol or acetone.

### Background of the Invention

With the introduction of new and more durable types of synthetic resins and polymers in protective and decorative coatings, the problem of stripping has become increasingly difficult. Prior art paint removers typically contain blends of aromatic solvents, ketones, paraffin, methanol and diluents, for example, mineral spirits in order to reduce raw material costs. Those prior art strippers made with the foregoing solvents focused industry attention on the flammable and toxic characteristics.

As an alternative, improved prior art solvents utilize chlorinated solvents as a way to extend the usefulness of paint strippers while simultaneously reducing fire hazards associated with their use. Methylene chloride (dichloromethane) has been almost universally used as the solvent of choice, although some compositions have been formulated with ethylene dichloride and trichlorethylene, among other chlorinated hydrocarbon solvents. Many of the chlorinated hydrocarbons including methylene chloride are suspected carcinogens and methylene chloride is designated a hazardous substance under the Federal Hazardous Substance Act (FHSA). In addition, methylene chloride, as a high vapor pressure chlorinated solvent, probably contributes to atmospheric ozone depletion. Methanol is a poison if ingested and the U.S. Consumer Products Safety Commission requires a warning designating it as a poison when its concentration in consumer products exceeds 4 percent by weight. Toluene contains trace quantities of benzene, a known carcinogen, and acetone with its high vapor pressure and low flash point is highly flammable in many prior art compositions.

Paint strippers which include methylene chloride can be formulated to soften, lift and blister paint films. The stripper which is applied to the substrate by brush, immersion

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or spray and allowed to loosen the paint or varnish has an available work life of only several minutes because of the tendency of the methylene chloride to evaporate rapidly. Because of the high vapor pressure of methylene chloride, in an effort to extend the work life of the stripper made with that solvent, paraffin is often included as an evaporation suppressant. The paraffin functions by forming a thin film at the surface of the paint remover which retards evaporation. This thin film formation and resultant evaporation suppressant activity of paraffin is produced by the evaporation and surface chilling which occur when methylene chloride or mixtures containing methylene chloride are exposed to air. However, methylene chloride strippers must be handled carefully since they are highly toxic and can cause severe skin irritation. Further, even with the incorporation of paraffin, the volatility of the methylene chloride is still very high and when methylene chloride is formulated with, for example, toluene, the resulting high levels of hydrocarbon emissions necessitate the provision of good ventilation to safely use the methylene chloride paint strippers.

A further disadvantage of paint strippers formulated with methylene chloride is that the short work life compromises the ability of the stripper to function efficiently in many applications. For example, many industrial uses of paint strippers involve stripping accumulated paint from equipment and walls. Paint strippers utilizing methylene chloride, suffer from the disadvantage that the short work life prevents the stripper from entering crevices in the accumulated paint layers thus creating the necessity, in some cases, of applying a large number of applications of stripper to remove a number of accumulated paint layers. Furthermore, the short work life of methylene chloride creates a recoating phenomenon in which stripped paint recoats the surface of one of the accumulated paint layers or the surface to be stripped as the solvent evaporates.

To remove paint on vertical surfaces, a thickener is generally added to avoid the problem of stripper flowing off the surface before it has had a chance to remove the paint layer. It is preferable to add a thickener, which in combination with the other stripper ingredients, may be easily washed

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off the surface after the stripping has taken place. However, if a paint stripper is too thick, it may be used on vertical surfaces, but it may not be amenable to spraying. Furthermore, the use of methylene chloride within the stripper may evaporate, thus producing a stripper which varies greatly in viscosity as a function of the evaporation time of the solvents in the stripper. Depending upon the required viscosity and the compatibility of the paint stripper ingredients with the paint stripper thickening system it may be necessary to reformulate the paint stripper with each application. Because of their high flash point, stripping compositions of the present invention may be adapted for a number of applications by simply changing the amount and type of thickener employed without modifying the other components. In certain preferred embodiments of the present invention, organoclay rheological additives, especially including the montmorillonite clays such as stearalkonium hectorite are utilized to produce paint stripper compositions exhibiting enhanced characteristics. Surprisingly, the use of organoclay rheological additives produces highly active paint stripping compositions which exhibit excellent activity even when up to about 45% by weight water is included in the formulation.

#### Objects of the Present Invention

It is an object of the present invention to provide paint stripper compositions which are safe and effective to remove paint and other polymeric coatings in protective and decorative coatings on flexible and inflexible surfaces.

It is also an object of the present invention to provide compositions which are non-flammable and effective paint strippers which do not contain petroleum based aromatic compounds.

It is an additional object of the present invention to provide stripping compositions and methods for removing conventional lacquers, varnishes, enamels, for example, drying oil type, alkyds, or drying oil-alkyd modified enamels, epoxy esters, epoxy amides, amine-catalyzed epoxies, acrylics and polyurethanes, among others.

It is a further object of the present invention to provide a method of stripping polymeric coatings using the compositions of the present invention.

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It is yet another object of the present invention to provide a method of making the compositions of the present invention, especially those compositions which are thixotropic and utilize organoclay rheological additives and gel formation.

It is yet an additional object of the present invention to provide paint stripper pastes comprising organoclay thickeners in combination with other stripping components of the present invention.

It is still a further object of the present invention to provide efficient paint stripping compositions which are pleasant, non-offensive in odor and which do not emit harmful fumes.

It is yet an additional object of the present invention to provide an effective paint and varnish removing composition which is non-caustic, has a neutral pH and will not harm wood.

Still an additional object of the present invention is a stripping composition that stays wet and active longer, may cover more surface area than prior art compositions and strips through multiple layers of paint and varnish.

#### Summary of the Invention

These and other objects of the present invention are accomplished by paint stripper compositions having high flash points, for example, above about 120°F and an absence of low-boiling VOC's comprising:

- a. about 0% to about 60% by weight of a terpene compound containing at least 10 carbon atoms;
- b. about 10% to about 69% by weight of N-methyl pyrrolidone or its equivalent; and
- c. about 0% to about 20% by weight of an emulsifying surfactant.

Compositions of the present invention may further comprise one or more of the following: about 5% to about 15% by weight of a solvent extender or bulking agent, about 5% to about 60% of a high flash point, high  $K_B$  (kauri-butanol) solvent selected from the group consisting of ethylene carbonate, propylene carbonate, butylene carbonate, glycols and glycol ethers, about 0% to about 45% water, from 0% to about 8% by weight of a thixotropic or thickening agent, in certain cases

*significantly enhancing the desired asphaltibility and*

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preferably an organoclay thickening agent, for example, stearalkonium hectorite, about 0.25% to about 20% by weight of an odor masking component, for example d'limonene and mixtures, thereof. In compositions containing organoclay rheological additives, about 10% to about 45%, preferably about 30 to 45% of a polar solvent, especially including water, ethanol and mixtures of water and ethanol is included in combination with the organoclay rheological additive. Depending upon the desired pH of the compositions, a base or acid may additionally be added to adjust the pH of the compositions of the present invention. In addition, for certain industrial applications it has surprisingly been found that the use of certain halogenated hydrocarbons containing both chlorine and fluorine in an amount ranging from about 2% up to about 40% results in compositions exhibiting especially advantageous stripping efficiencies.

The paint stripper compositions of the present invention are safe, effective and can be used in a wide variety of applications. Environmentally safe, biodegradable compositions of the present invention solve many of the problems associated with the use of low-boiling chlorinated hydrocarbons in prior art paint strippers, including the health hazards and environmental problems associated with the use of those strippers. The compositions of the present invention have a high flash point, preferably above about 93°C (200°F), are generally biodegradable and can be formulated to be highly active in the pH range of about 6.0 to about 8.0, a particularly useful advantage over prior art compositions. In addition, certain embodiments of the present invention provide the advantage that they can be used to treat a coated surface and then removed by simply washing the treated surface with water.

In one particularly preferred aspect of the present invention, stripper compositions of the present invention are adsorbed onto paper or fabric sheets and used to strip paint and other coatings from a surface. Sheets to which are adsorbed the compositions of the present invention are very easy to handle and provide simple and efficient means to remove paint from large surface areas, for example walls, doors and panels.

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In another preferred embodiment of the present invention, the thixotropic agent is an organoclay rheological additive and the compositions are used as pastes. In these compositions the amount of polar solvent that can be employed without impacting the activity of the paint stripper composition may be as high as about 45% by weight. In general, water is used as the polar solvent, but preferred compositions include mixtures of ethanol and water. In addition, other polar solvents, for example, isopropanol, methanol, acetone and methylethyl ketone alone or in combination may also be used, but these are less preferred than is a mixture of ethanol and water.

The compositions of the present invention can be used to strip or remove conventional lacquers, varnishes, enamels, epoxy esters, epoxy amides and acrylics, among other polymeric coatings. The time required to strip paint finishes completely is competitive with that of conventional strippers containing chlorinated solvents, and ranges from about 5 minutes to about 80 minutes, with a preferred range of about 10 to about 35 minutes. In the case of organoclay containing paste compositions, the time required to strip paint may be as short as 5 to 10 minutes. Certain embodiments of the present invention may be used to strip multiple layers of coatings off of surfaces by exposing the surface to the paste compositions of the present invention for periods ranging from 15 minutes to more than 24 hours, sometimes even several days.

In removing coatings from surfaces, the compositions of the present invention produce a wrinkling or blistering of the polymer coating. While not to be limited by way of theory, this fundamental blistering is believed to be produced by swollen gel. When normally insoluble high polymers are placed in contact with an active solvent, the polymer absorbs the solvent and expands until an equilibrium condition occurs. In order to accommodate the increased volume and still remain within the original surface perimeter, the polymer tends to fold or wrinkle so as to relieve internal stresses. The solvent mixtures solvate the active groups of the polymer, and in so doing, reduce solid to solid adhesion. The forces which build up in the polymer generally, as a result of a combination of swelling and weakened adhesion, are sufficiently high

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to tear the film loose from the surface. Wrinkle formation occurs because solvent penetration into the polymer and deterioration of the polymer cohesive bonds proceed at approximately the same speed. There is rapid parting of the film from the substrate and almost as rapid rupturing of the film itself.

In the case of organoclay containing compositions, it is believed that enhanced activity associated with these compositions may be the result of water and other polar solvents in association with the organoclay rheological additives producing a solvent effect in which the terpenoids, N-Methylpyrrolidone or its equivalents and the other active components in the compositions exhibit enhanced interaction with the polymeric coating to be stripped.

In the method aspects of the present invention, compositions of the present invention are used to strip and/or remove polymeric coatings on a number of surfaces. In a particularly preferred method of using the compositions of the present invention, stripping composition preferably adsorbed to fabric or paper is placed in contact with a coated surface and after a period of time sufficient to soften and/or blister the underlying coating, the paper is removed along with the softened coating in one easy step.

Depending upon the formulation of the composition to be used, further method aspects of the present invention include making the compositions of the present invention using a step-wise procedure in which thixotropic agents are mixed separately from the active ingredients. Certain thixotropic compositions, for example methylcellulose, such as Methocel 311, available from Dow Chemical Corp. Midland, Michigan, hydroxypropyl cellulose, Klucell™, available from Hercules, Inc., Wilmington, Del., ethyl cellulose, such as Bermocol PR™, available from Seaboard Sales, New Jersey) and organoclays modified with quaternary compounds (high medium and low polar), such as Bentone 27 and stearalkonium hectorites, as well as other Bentones including Bentone EW™ and Bentone LT™, products of NL Industries, New Jersey, USA are included in amounts ranging from 0% to about 8%, preferably about 0.02% to about 5%, and most preferably about 1 to about 3% of thixotropic agent. Gel formation which occurs separate

from the other active agents in the compositions of the present invention produces a composition exhibiting acceptable thixotropy for stripping or removing polymeric coatings on vertical surfaces and enhanced stripping activity. In addition, this method aspect of the present invention results in a composition that is much easier to make than compositions in which gel formation occurs in the presence of the active ingredients and the quality control of which can be easily monitored.

#### Detailed Description of the Invention

The compositions of the present invention comprise from about 0% to about 60% by weight of a terpene compound. As used herein, the term terpene refers to a class of acyclic and cyclic unsaturated compounds derived from natural essential oils and resins having at least 10 carbon atoms. Terpenes that find use in the present invention include alcohols and aldehydes as well as unsaturated hydrocarbons. Any number of terpene compounds, including combinations of these terpenes may be used in the present invention as the base or primary solvent as polymer absorbents. Preferred terpene compounds are those compounds which have a  $K_B$  value of at least about 108 (the same value as toluol, an aromatic hydrocarbon which is the solvent of choice in many prior art strippers) and a flash point above about 120°F.

In the case of organoclay containing compositions which generally include about 10% to about 45%, preferably 30% to about 45% of a polar solvent including water, the amount of terpene included generally comprises about 0% to about 50% by weight of the composition.

$K_B$  is a measure of the solvency of a hydrocarbon. In general, the higher the  $K_B$  value, the greater the general solvent power of the hydrocarbon under test conditions described by ASTM D1133. To determine  $K_B$  value, a hydrocarbon sample is added to a standard solution of kauri gum in butyl alcohol until sufficient kauri gum precipitates to blur vision of 10 point type viewed through the flask. When used in varnish, lacquer and enamel formulations, a hydrocarbon diluent with a high  $K_B$  value dissolves relatively large quantities of solids.

The preferred terpene compounds for use in the present invention are represented by unsaturated hydrocarbons,

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alcohols and aldehydes having at least 10 carbon atoms and include 3,7-dimethyloctanol, alpha-pinene, beta-pinene, delta-3-carene, citronellal, citronellol, hydroxycitronellal, d-limonene, linalool, gamma-terpinene, tetrahydrolinalool and terpineol, among others, with cyclic terpenes being preferred. Especially preferred terpenes for use in compositions of the present invention include dipentene, because of its high  $K_B$  value.

Compositions of the present invention comprise about 0% to about 60% by weight of a terpene compound. Preferred compositions of the present invention may comprise between about 15% to about 50% by weight of a terpene compound, and most preferably, comprise between about 15% to about 30% by weight of a terpene compound. Organoclay containing compositions according to the present invention generally comprise about 0% to about 50% by weight of a terpene, preferably about 10% to about 25% by weight terpenes and most preferably about 15% by weight terpenes. The terpene compounds of the present invention are used in the above-defined weight ranges because these weight ranges are effective to cause polymer expansion which facilitates paint stripping.

In certain embodiments where the preferred terpene, dipentene is used, a malodor associated with thiols sometimes occurs. In such cases, a natural odor masker, for example, d'limonene, among other odor masking agents, may be added in an amount ranging from about 0.25% to about 20% to make the compositions more pleasant in odor.

*NMP*

Compositions of the present invention also comprise between about 10% to about 69% by weight of N-methylpyrrolidone or its equivalent. N-methylpyrrolidone is a highly polar, water-miscible solvent which is also miscible with almost all conventional organic solvents. It is also soluble in a wide range of polymers used in the coatings industry. It serves to dissociate chemical groupings in the coatings which are not dissociated by the primary terpene solvent. Compositions containing as little as 10% by weight N-methylpyrrolidone are useful, but are slower in stripping efficiency than compositions containing higher weight percentages of N-methylpyrrolidone. Other solvents similar to N-methylpyrrolidone in solubility and solvent power may be used

*200-63**promise (an und 20%)*

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in the present invention, and may include structural analogues of N-methylpyrrolidone, for example N-ethylpyrrolidone and N-propylpyrrolidone, among other solvents. Of course, it will be recognized by those of ordinary skill in the art that a number of variations may be made to the structure of N-methylpyrrolidone without significantly affecting the physicochemical properties of the underlying stripping composition. Where required, one of ordinary skill in the art will know to vary the amounts and types of the other components within the stripping composition to produce an efficient paint stripping composition.

*UAP* ( In organoclay containing compositions according to the present invention, the amount of N-Methylpyrrolidone may comprise as much as about 69% by weight, but preferably comprises about 20% to about 40% by weight and most preferably about 25 to about 30% by weight.

The present invention utilizes pyrrolidone compounds such as N-Methylpyrrolidone and similar compounds having a lactam moiety, but clearly N-methylpyrrolidone is preferred. As described hereinabove, the nitrogen of the pyrrolidone ring may be unsubstituted or substituted with alkyl groups. However, as the number of carbon atoms within the pyrrolidone compound increases beyond 3 (propyl), the water-miscibility decreases, thus affecting the ability of stripping compositions to accommodate water for microgel formation or for removal of the stripping composition from surfaces. In addition, such substitution may affect the solvent power of the pyrrolidone compound used, thus necessitating adjustments in the other components of the stripping composition. In addition to nitrogen substitution, the carbon atoms of the pyrrolidone ring may also be substituted, but such compositions for use in the present invention are less preferred than N-Methylpyrrolidone because of the cost of such compositions and the reduced polarity that often results from such substitution. For example, N,5-dimethyl 2-pyrrolidone, 2-pyrrolidone, 3,3-dimethyl 2-pyrrolidone may be used in the present invention, but are less preferred than is N-Methylpyrrolidone.

The 2-keto group of N-methylpyrrolidone is not a requirement for the pyrrolidone compound of the present invention and certain pyrrolidine analogs, for example, N,2-

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dimethylpyrrolidine, 2,4-dimethylpyrrolidine, 2,5-dimethylpyrrolidine and N-methylpyrrolidine, among others may also be used in the present invention. Pyrrolidone and pyrrolidine compounds which may be used in the present invention may be less polar than N-Methylpyrrolidone thus necessitating adjustment to the amount and type of surfactant and/or terpene compound used.

The terpenes and N-methylpyrrolidone are miscible, and N-methylpyrrolidone itself is miscible in water. However, the terpenes and water are not miscible. Therefore, in order to provide compositions which have sufficient quantities of terpene compounds to solubilize and expand coating polymers yet accommodate minor amounts of water for microgel formation and greater amounts of water for ease of removal, one or more emulsifying surfactants are included in compositions of the present invention in an amount equal to about 0% to about 20% by weight and preferably about 2% to about 15% by weight of the paint stripping compositions of the present invention. In the organoclay containing compositions according to the present invention, when they are included, emulsifying surfactants generally comprise about 1% to about 10% by weight, preferably about 2% to about 7% by weight and most preferably about 3% to about 5% by weight.

*Surfactants* Numerous surfactants capable of emulsifying the terpenes and other components of the present invention may be employed, including but not limited to linear or branched chain alcoholic ethoxylates and ethoxysulfates, alcohol ethoxylates, polysorbate esters, ethoxylated alkylphenols, for example, polyethoxynonylphenols, phenoxy polyalkoxy alcohols, for example, nonylphenoxypoly(ethyleneoxy)ethanol and nonylphenoxypolyethoxyethanol, alkyl and alkyl ether sulfates and sulfonates, for example, dodecylbenzenesulfonic acid, alkyl and dialkyl succinate compounds, phosphate esters, for example phosphate esters of long-chain alcohol ethoxylates and combinations of these surfactants. Particularly preferred surfactants for use in compositions of the present invention include the phosphate ester surfactants, for example PD-600<sup>TM</sup>, an alkaline stable mixture of mono and di-substituted phosphate esters of decylalcoholethoxylate, available from Chemax, Inc. (Greenville, South Carolina) and the Tryfact<sup>TM</sup>

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phosphate esters, a series of phosphate ester surfactants as the free acids or various salts, available from Emery Chemicals, Mauldin, South Carolina. Other preferred surfactants include polyoxyethylenenonylphenol (NP-6<sup>TM</sup> and NP-9<sup>TM</sup>, available from Chemax, Inc., Greenville, South Carolina) as well as the nonionic alkylphenoxy polyoxyethylene alcohols, the Makon<sup>TM</sup> series, available from Stepan Company, Northfield, Illinois. Alkylamine dodecylbenzenesulfonate, Ninate 411<sup>TM</sup>, available from Stepan Company, the linear alkylbenzene sulfonic acid surfactant Bio-soft s-100<sup>TM</sup>, available from Stepan, sodium 1-octane sulfonate, Bio-terge pas-8s<sup>TM</sup>, available from Stepan and Neodol 1-5<sup>TM</sup>, a nonionic surfactant having an average of 5.0 ethylene oxide units per molecule of alcohol, available from Chemax, Inc., are also useful in the present invention. A preferred surfactant for use in paint strippers having an alkaline or basic pH includes CSX-727, an alkaline stable surfactant also available from Chemax, Inc. These preferred surfactants may be used alone or in combination to enhance the activity of the terpinoid and N-methylpyrrolidone mixture and to produce paint strippers having excellent paint stripping activity.

The pH of the paint stripper compositions of the present invention may vary from highly acidic to highly alkaline within the range of about 2.0 to greater than 11.0. Preferred compositions generally have a neutral pH. Paint strippers of the present invention which are to be used as industrial or military paint strippers may be very alkaline, i.e., have a pH above about 10.0, because enhanced paint stripping activity has been exhibited by paint strippers having a pH above about 10.0. However, a number of preferred embodiments of the paint stripper of the present invention have a pH within the range of about 5.5 to about 8.0 and preferably within the range of about 6.0 to about 8.0. It is a particularly surprising result that paint stripper compositions of the present invention show substantial paint stripping activity in the industrial setting within a neutral pH range of about 6.0 to about 7.0. Thus, although an acidic or basic accelerator may be added to compositions of the present invention to increase stripping efficiency and shorten stripping time, such an acidic or basic accelerator is not necessary. In compositions

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to be used in the home by the consuming public, it is preferred to exclude an acidic or basic accelerator which generally reduces the safety of the compositions. In the preferred organoclay containing compositions, enhanced paint stripping activity is associated with a neutral pH, i.e., about 6.5 to about 7.5. This is a surprising result that such enhanced activity may be obtained without very high or very low pH conditions.

Although anionic, nonionic and cationic surfactants may be employed in certain embodiments of the present invention, it is preferable to employ nonionic surfactants. It may be preferable in certain embodiments of the present invention to include a mixture of surfactants to enhance the solubility of certain active components in oil or water. A surfactant mixture may serve to promote the stripping of the polymeric coating by the active components and enhance the removal of the stripper and stripped coating by a washing step.

Compositions of the present invention may also include a solvent extender in an amount equal to about 5 to about 15 percent by weight of the compositions. The solvent extenders in the present invention function primarily as bulking agents, but they may also function to enhance certain solubility characteristics of the compositions. In addition, the solvent extenders used in the present invention may be useful in certain embodiments for forming gel from thixotropic agents such as cellulose ethers used in certain embodiments of the present invention. Typical solvent extenders for use in the present invention include small chain alcohols, for example, ethanol and isopropanol, among others.

Organoclay containing compositions according to the present invention generally contain a polar solvent in an amount ranging from about 10% to about 45% by weight and preferably about 30% to about 45% by weight. Although higher weight percentages of polar solvent may be used, they are generally less preferred. A polar solvent for purposes of the present invention includes those solvents and mixtures of solvents which, in combination with sufficient amounts of organoclays such as, for example, the Bentones<sup>TM</sup>, produce a thixotropic gel after mixing under shearing force. Exemplary polar solvents for use in the present invention include water,

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ethanol, isopropanol, methanol, acetone and methylethyl ketone, among others, including mixtures of these solvents. Water, ethanol and mixtures of water and ethanol are clearly preferred.

Paint stripper compositions of the present invention may also include about 5% to about 60%, preferably about 20% to about 40% of a high flash point, high  $K_B$  (kauri-butanol) solvent selected from the group consisting of polycarbonates, including ethylene carbonate, propylene carbonate and butylene carbonate, and glycols and glycol ethers. This high flash point, high  $K_B$  solvent is added to increase the flash point of compositions as well as maintain the ability of the compositions to blister and remove polymeric coatings from surfaces. The paint stripping compositions of the present invention have a flash point of at least about 150°F, preferably at least about 175°F and most preferably at least about 200°F, flash points which are obtainable, at least in part by virtue of the amount of high flash point solvent added to the compositions. Preferred high flash point solvents for use in the present invention have a flash point of at least about 80°C and a  $K_B$  of at least about 300. In the organoclay compositions according to the present invention, the high  $K_B$  solvent preferably comprises about 5% to about 40% by weight of the compositions.

Glycols useful in embodiments of the present invention include ethylene glycol, propylene glycol and butylene glycol, among others. Glycol ethers may be selected from among a number of agents including ethyleneglycolmonomethylether, diethyleneglycolmonomethylether, triethyleneglycolmonomethylether, ethyleneglycolmonoethyl ether, diethyleneglycolmonoethyl ether, triethyleneglycolmonoethyl ether, ethyleneglycolmonopropylether, diethyleneglycolmonopropylether, ethyleneglycolmonobutylether, diethyleneglycolmonobutylether, triethyleneglycolmonobutylether, ethyleneglycolmonohexylether, diethyleneglycolmonohexylether, ethoxypropyleneglycolmonobutylether, propyleneglycolmonomethylether, di-propyleneglycolmonomethylether, tri-propyleneglycolmonomethylether, propyleneglycolmonopropylether, di-propyleneglycolmonopropylether, propyleneglycolmono-t-butylether, propyleneglycol-

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monobutylether and dipropylene glycolmonobutylether, among others. Preferably, propylene carbonate is used in compositions of the present invention because of its favorable flash point and  $K_B$  value as well as its general safety and lack of toxicity.

*edw*

Another component which may be added to the compositions of the present invention includes water in an amount equal to about 0% to about 45%. Preferably water comprises about 0.5% to about 20% by weight of the composition in those embodiments which do not contain organoclay rheological additives. In those compositions containing the organoclay rheological additives, for example, the stearalkonium hectorites such as the Bentonest<sup>TM</sup>, the amount of polar solvent including water comprises about 10% to about 45%, preferably about 30% to about 45% by weight of the compositions.

Water is added in certain embodiments of the present invention to provide solubility characteristics so that the stripping compositions, after they have blistered the paint or varnish coating a surface to be stripped, may be washed off from the surface with a stream of water. In addition, in certain embodiments containing a thixotropic agent such as the celluloses, water is added to aid the formation of gel. Of course, as the percent by weight of water changes, the other components of the compositions and in particular, the terpene and terpene emulsifying surfactant components may have to be adjusted to accommodate the additional water.

Compositions of the present invention may also include a thixotropic agent to increase viscosity so that compositions of the present invention may be applied to vertical surfaces without running off. Compositions of the present invention may comprise between 0% and about 8% of a thixotropic agent, preferably about 0.02% to about 5% and most preferably, about 1% to about 3% by weight of a thixotropic agent. Certain thixotropic compositions, for example methylcellulose, such as Methocel 311<sup>TM</sup>, available from Dow Chemical Corp., Midland, Michigan, hydroxypropyl cellulose, Klucell<sup>TM</sup>, available from Hercules, Inc., Wilmington, Del., ethyl cellulose ethers, such as Bermocol PR<sup>TM</sup>, available from Seaboard Sales, New Jersey) and organoclays modified with quaternary compounds (high, medium and low polar), such as the Bentonest<sup>TM</sup>, for example,

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Bentone 27, Bentone EW<sup>TM</sup> and Bentone LT<sup>TM</sup>, products of NL Industries, New Jersey, USA are preferred, with the Bentones<sup>TM</sup>, especially Bentone EW<sup>TM</sup> being particularly preferred.

The organoclays additives are preferably utilized as thixotropic agents in the varnishes and paint remover compositions of the present invention. The organoclays which can be utilized in the present invention generally depend on wetting, shear and polar activation for rheological development. One class of the organoclays, the high performance hydrophilic organoclays, including Bentone EW<sup>TM</sup> and Bentone LT<sup>TM</sup>, depend only upon shear, hydration and wetting for activity. These hydrophilic organoclays offer the formulator opportunities to incorporate large quantities of water, i.e., up to 45% and higher by weight in paint remover compositions thereby reducing cost and eliminating hazards associated with flammable compositions without reducing the efficacy of the compositions.

Compositions of the present invention include a thixotropic agent to increase the viscosity of the compositions to a level which will result in a paint stripper which will remain on vertical surfaces for a period of time at least long enough to blister the polymeric coating on such surface and exhibit efficient stripping efficiency. Preferred thixotropic agents include ethyl cellulose and Bentone EW<sup>TM</sup>, among other agents, which produce a gel when mixed into the paint stripping compositions of the present invention.

Gel formation may be produced by any method including simply mixing the thixotropic agent into the composition at any time. However, it is advantageous to form a gel separately from the active components for ease of formulation. Gel formation which occurs separately from the other active agents in the compositions of the present invention produces a composition exhibiting acceptable thixotropy for stripping or removing polymeric coatings on vertical surfaces and enhanced stripping activity. In addition, this method of the present invention comprising forming gel separately from the active agents (in the presence of water and/or a bulking agent such as ethanol) and then adding the microgel mixture to the separately mixed active component results in a composition that is

much easier to make and the quality control of which can be easily monitored. The texture of these compositions made by this method is often superior as well.

In certain embodiments of the present invention, especially those compositions which may be used to remove especially difficult polymers, for example in many military and industrial applications, compositions of the present invention, may additionally comprise between 0% and 40% by weight of a chlorinated, fluorinated hydrocarbon (chlorofluorocarbons, CFCs or HCFCs) having a boiling point of at least about 45°C. The weight percent of CFCs used in compositions of this aspect of the present invention ranges between 0% and 40%, preferably about 1% to about 25% and most preferably about 2% to about 18% by weight of the paint stripping compositions. The CFCs are added to compositions of the present invention to provide an additional component to blister the polymeric coatings on surfaces. Paint stripping compositions of the present invention which include CFCs generally exhibit greater stripping efficiencies than those which do not. Surprisingly, certain organoclay compositions according to the present invention which do not include CFCs (HCFCs) are active against especially difficult coatings including cured coatings on military hardware.

A number of CFCs may be used in the present invention, but preferred CFCs have a boiling point that exceeds at least about 45°C. Although lower boiling point CFCs may be used, they are difficult to work with because of their tendency to volatilize rapidly. CFCs used in the present invention are non-flammable and do not flash and are advantageously included in certain compositions of the present invention to modify the stripping efficiency as well as increase the flash point of the composition. Preferred CFCs include those in which the number of fluorine atoms is the same or greater than the number of chlorine atoms and the number of carbon atoms ranges from 2 to 4. Exemplary CFCs include  $\text{CCl}_2\text{F}_2$  (CFC 113<sup>TM</sup>, available from DuPont, Wilmington Delaware),  $\text{CHCl}_2\text{CF}_3$  (CFC 123<sup>TM</sup>, also available from DuPont),  $\text{CHClF}_2\text{CH}_2\text{Cl}$  (1,2-dichloro-1,1-difluoroethane),  $\text{CHClFCH}_2\text{Cl}$  (1,2-dichloro-1-fluoroethane),  $\text{CCl}_2\text{FCCl}_2\text{F}$  (1,1,2,2-tetrachloro-1,2-difluoroethane),  $\text{CHClFCHCl}_2$  (1,2,2-trichloro-1-fluoro-ethane),  $\text{FCl}_2\text{CCl}_3$

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(fluoropentachloroethane),  $F(CH_2)_3Cl$  (1-chloro-3-fluoropropane),  $CH_2ClCF_2CH_3$  (1-chloro-2,2-difluoropropane),  $CH_2ClCClFCH_3$  (1,2-dichloro-2-fluoropropane),  $CF_2ClCClFCClF_2$  (1,2,3-trichloro-1,1,2,3,3-pentafluoropropane),  $CF_2ClCCl_2CF_2Cl$  (1,2,2,3-tetrachloro-1,1,3,3-tetrafluoropropane) and  $Cl(CH_2)_4F$  (1-chloro-4-fluoro-butane), among others. Most preferably, HCFC 123<sub>tm</sub> is used in certain embodiments of the present invention in amounts ranging from about 2% to about 40%, preferably about 2% to about 25% and most preferably about 2% to about 18% by weight of the compositions.

The compositions of the present invention are generally applied to the surface to be stripped in the usual manner, i.e., the compositions are first applied by brush, spatula or other applicator and then are applied to the surface to be stripped. Alternatively, the compositions may be sprayed onto a surface in a continuous flow method which recirculates the composition. Generally, the compositions are in contact with the surface for a period of time sufficient to produce a blistering of the polymeric coating, after which time, the blistered coating is removed by contacting the coating with an abrasive pad. Alternatively, the coating is removed by simply lifting the blistered coating off of the surface or by spraying the blistered coating with water. Certain embodiments of the present invention, including the impregnated fabric and the organoclay containing compositions may be applied to a coated surface for periods exceeding 24 hours.

In a particularly preferred embodiment of the present invention, the stripping compositions of the present invention are adsorbed onto a fabric piece comprising cloth or paper and applied to a surface to be stripped. Generally, a non-woven, highly absorbent fabric, without chemical binders is used. Although any fabric or paper which absorbs the compositions of the present invention and is stable to the compositions of the present invention may be used in this aspect of the present invention, preferred fabrics include a thermally bonded composite blend of cellulose and propylene (cellulose/propylene), typically a 50-50 percent by weight blend of each component. Additionally, different fabrics based upon polyester, rayon/polyester blends and woodpulp/polyester blends (the

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Sontara<sup>TM</sup> fabrics from DuPont), among others may be used efficiently. The fabric may range in thickness and abrasive quality, but typically, preferred fabrics range in thickness from about 0.28 mm up to 1.02 mm.

Generally, cloth having a basis weight of approximately 2 ounces/yard is satisfactory for use in the present invention, but the basis weight may vary without impacting the stripping efficiency of the compositions of the present invention. Fabric to which compositions of the present invention are adsorbed are generally packaged in a barrier bag to prevent the transmission of vapor. Generally, the barrier bag is made of aluminum foil or other material which will not transmit vapor from the paint stripping compositions of the present invention.

The procedure for making the fabric applicator of the present invention is quite simple. In general, fabric is cut or slit to the appropriate size depending upon application, folded and inserted into the barrier bag which is sealed on three of four sides. Thereafter, a metered volume of stripper composition is injected into the open end of the barrier bag which is then immediately sealed. After the injection of paint stripper and during storage, the paint stripper composition impregnates the fabric applicator.

The fabric applicator is designed to blister and remove coatings from surfaces. It is especially useful when applied to vertical surfaces because it adheres when applied. The fabric applicator and paint stripper composition is designed so that there is sufficient tension between the applicator and the applied surface to retain it on the surface of the wall. By design, the applicator continuously applies composition to the surface to be stripped until the paint has lifted. A particularly advantageous feature of this aspect of the present invention is the fact that paint stripper composition will not evaporate from the applied surface because the fabric significantly reduces evaporation. Thus, the fabric embodiment of the present invention may be used on particularly difficult coatings and on multiple layers where the contact time between the paint stripper and the coated surface needs to be longer than in less rigorous applications.

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To use the fabric aspect of the present invention, the fabric applicator to which stripper composition is adsorbed is placed on a coated surface. The applicator remains in contact with the coated surface for a period sufficient to blister the underlying coating. Such period may be 24 hours or longer for particularly difficult coatings. Once the coating blisters, the applicator is simply peeled off, usually with the underlying coating fixed to the applicator. Often there is no need to scrape or abrade the coating.

The following characteristics are identifiable in compositions of the present invention:

- (1) Increased stripping efficiency, i.e., reduced stripping time, and increased penetration through multiple layers comparable to commercially available methylene chloride strippers.
- (2) Raised flash point of the formulations to provide a stripper which will not evaporate within a stripping period.
- (3) Minimized objectionable odors.
- (4) Neutral pH for widest possible range of utilization and safety.
- (5) Highest possible TLV (threshold limit value) and exposure, lowest toxicity, and maximum environmental acceptability.
- (6) A water soluble stripper composition which can be used in industrial settings where removal is performed by spraying a stripped surface with water.
- (7) Impregnate absorbent material for application of impregnated sheets to vertical surfaces to be stripped.
- (8) Fabric applicator utilizes absorbent materials with properties of low density, high absorbancy and durability, without binders, and non-effected by ingredients of composition.

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(9) Organoclay containing compositions containing high concentrations of water exhibiting enhanced stripping activity.

The following examples are provided to illustrate the present invention and should not be misunderstood to limit the scope of the present invention in any way.

### EXAMPLES

#### General Procedure for Formulating Examples 1-22:

Each component of the paint stripping composition of the present invention is weighed out. Generally, the terpene and N-methylpyrrolidone or its equivalent are first blended. A high flash point, high  $K_B$  solvent may then be added to the above mixture. The surfactant system is first mixed (surfactant and water) and then added to water and mixed. The water/surfactant mixture is then added to the terpene and N-methylpyrrolidone. In cases where a thixotropic agent was added, thixotropic agent was added to water and a bulking agent which are blended together using an air-driven agitator until homogeneous. The mixture containing thixotropic agent is then added to the terpene/N-methylpyrrolidone/surfactant mixture of components. This mixture is then agitated until the composition is homogeneous.

In the case of organoclay rheological additives, the clay is first mixed with the polar solvent, generally a mixture of ethanol and water to produce phase A. Phase B, which contains the other ingredients, is prepared separately. The ingredients comprising phase B may be added in any order. After mixing, phase B is added to phase A and subjected to vigorous mixing under shearing force to produce a uniform gel-like or paste-like product. The composition is then tested for stripping activity.

#### General Testing Procedures

Standardized test panels coated with the substrate to be stripped are prepared. Enamels, alkyds, epoxies, latex, acrylics and conventional lacquers were tested. Testing was performed on metal plates with three coats of paint, each coat of different color so as to permit observation and time

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measurement of coating penetration. In the case of organoclay containing compositions, the testing was performed on a number of metal plates as described above, wood panels, boards and sections of doors. Stripping efficiency is defined as the ratio of the time required to observe blistering divided by 20 (minutes).

The testing of strippers occurred in three steps. In the first step, utilizing standard test plates, a droplet approximately 1/2 inch in diameter of a stripping composition is placed onto the test plate, and the time until the coating blisters is then measured. Compositions which showed a stripping time of less than 40 minutes are then tested in a second test. In the second test, the coating composition is then applied onto a substrate by brushing and the time to blister is measured. In a third test, candidate compositions are tested for stripping efficiency, odor, ease of use and product satisfaction. In this third test, two methods of stripper application have been utilized. The first method is continuous flow, where the composition is applied to the substrate by pumping (air pressure) and recirculation of the solution while an abrasive pad is used to remove the coating. This application requires a liquid solution in order to facilitate flow and pumping. In a second method of the third test, a thixotropic solution (to minimize flow and dripping when applied) is brushed onto irregular, ornate or vertical surfaces. This test is utilized to determine the efficacy of thixotropic and especially organoclay containing compositions according to the present invention. In the third test, the formulations of the present invention were compared to conventional paint strippers. The most challenging substrates were chosen to be stripped, to establish the performance criteria expected of competitive paint strippers.

EXAMPLE 1  
Alkaline Stripper System #1

COMPONENT	WEIGHT PERCENT
DIPENTENE	47
ISOPROPANOL	6
N-METHYLPYRROLIDONE	33

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SURFACTANT SYSTEM (WITCOLATE D5-10) (NP-9)	8
WATER/KOH	4
	2

Procedure: Dipentene, isopropanol and N-methylpyrrolidone were mixed separately. Surfactant system and water were first mixed separately and then added to above mixture.

Result: Useful compositions for stripping. Stripping time was about 60 minutes. Phase separations and solution stratification were evidenced.

**EXAMPLE 2.**  
Alkaline Stripper System #2

COMPONENT	WEIGHT PERCENT
DIPENTENE	47
ISOPROPANOL	6
M' PYRROLIDONE	33
SURFACTANT SYSTEM (WITCOLATE D5-10) NP-6 [polyoxyethylene (6) nonyl phenol]	8
WATER/KOH	4
	2

Procedure: Same as example 1.

Results: pH in the alkaline range. Stripping time was about 60 minutes. Useful stripper composition which evidenced phase-separation upon standing. Must be vigorously mixed before use.

**EXAMPLE 3**  
Alkaline Stripper System #3

COMPONENT	WEIGHT PERCENT
DIPENTENE	47
ISOPROPANOL	6
N-METHYLPYRROLIDONE	33
SURFACTANT SYSTEM: CSX-727/KOH (75% solution) KASIL	12
WATER	2

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Procedure: Same as examples 1 and 2.

Results: pH alkaline. Composition effective, but turbid. Stripping time was about 60 minutes. All components could not be completely solubilized.

**EXAMPLE 4**  
Alkaline Stripper System #4

COMPONENT	WEIGHT PERCENT
DIPENTENE	47
ISOPROPANOL	6
N-METHYL PYRROLIDONE	33
SURFACTANT SYSTEM: CSX-727/KOH (50%) KASIL#1	12
WATER	2

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Procedure: General method, as per examples 1-3

Results: Same as example 3- produced a useful but turbid suspension. Not all components solubilized.

**EXAMPLE #5**  
Alkaline Stripper System #5

COMPONENT	WEIGHT PERCENT
KOH SOLUTION (75%)	33
KABIL #1	32
CSX 727	5
METHYL PYRROLIDONE	10
FURFURYL ALCOHOL	20

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Procedure: Same as general method. KOH solution, CSX 727 and Kabil #1 mixed separately until completely homogeneous.

Results: pH is alkaline. Stripping efficiency marginal (50

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min. to soften - no lifting evident).

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EXAMPLE 6  
Alkaline Stripper System # 6

COMPONENT	WEIGHT PERCENT
KOH SOLUTION (75%)	20
KASIL #1	32
CSX 727	5
DIPENTENE	13
N-METHYL PYRROLIDONE	10
FURFURYL ALCOHOL	20

Procedure: General method, same as example 5.

Results: pH is alkaline, somewhat malodorous. Marginal stripping efficiency evidenced. Results similar to example 5. Malodor associated with dipentene at high pH. Alkaline based strippers (Examples 1-6) showed utility, but compositions needed to be buffered to produce a long shelf-life product.

EXAMPLE 7  
Neutral pH Stripper System #1

COMPONENT	WEIGHT PERCENT
PROPYLENE CARBONATE	45
N-METHYL PYRROLIDONE	25
PD-600 (phosphate ester)	7
GLIDCOL 95 (pine oil derivative with high K <sub>B</sub> )	3
DIPENTENE	7
ISOPROPANOL	9
WATER	3
METHYL CELLULOSE	1

Procedure: Follows general method. About half of water is used to create surfactant/water mixture which is added to mixture of propylene carbonate, n-methylpyrrolidone, glidcol 95 and dipentene. Other half of water is used with isopropanol and methyl cellulose to produce microgel mixture which is

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added to mixture of other components.

Results: Compositions showed better stripping efficiency than high pH systems. Stripping efficiency slightly reduced by methyl cellulose thickener.

**EXAMPLE 8**  
Neutral pH Stripper System #2

COMPONENTS	WEIGHT PERCENT
PROPYLENE CARBONATE	45
N-METHYL PYRROLIDONE	25
PD-600	7
GLIDSOL 150 (pine oil derivative with high $K_B$ )	3
DIPENTENE	7
ISOPROPANOL	9
WATER	3
METHYL CELLULOSE	1

Procedure: Same as Example 7.

Results: Composition showed results similar to Example 7.

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**EXAMPLE 9**  
**Neutral pH Stripper System #3**

COMPONENT	WEIGHT PERCENT
PROPYLENE CARBONATE	45
n-METHYL PYRROLIDONE	25
PD-600	7
GLIDSOL 100 (pine oil derivative with high $K_B$ )	3
DIPENTENE	7
ISOPROPANOL	9
Water	3
METHYL CELLULOSE	1

**Procedure:** Same as examples 7 and 8.

**Results:** Examples 7-9 utilize pine oil derivatives, which, although high in  $K_B$  value, with higher flash points and less odor than dipentene tend to inhibit stripping efficiency to a certain degree compared to dipentene. Dipentene exhibits superior substrate penetration of paints and varnishes to other terpenoid solvents tested.

**EXAMPLE 10**  
**Neutral pH Stripper System #4**

COMPONENT	WEIGHT PERCENT
PHENOXY ETHANOL (DIETHYLENE GLYCOL MENOPHENYL ETHER)	20
DIPENTENE	20
D'LIMONENE	10
N-METHYL PYRROLIDONE	15
PD-600	8
WATER	2

**Procedure:** Same as examples 1-6 and general method.

**Results:** Good solvent properties- good stripping efficiency, about 30 minutes. Thoroughly miscible - no objectionable odors. Flash point slightly too low.

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**EXAMPLE 11**  
Neutral pH Stripper System # 5

COMPONENT	WEIGHT PERCENT
PHENOXY ETHANOL (DIETHYLENE GLYCOL MONOPHENYL ETHER)	20
DIPENTENE	20
D'LIMONENE	10
N-METHYL PYRROLIDONE	40
PD-600	8
PROPYLENE CARBONATE/ETHYLENE CARBONATE (50/50)	25
WATER	2

Procedure: Same as example 10 and general method. Surfactant and water mixed separately and added to a mixture of the other components.

Results: Composition showed good stripping efficiency. But less favorable attributes of phenoxy ethanol from a safety standpoint associated with its use than for other surfactant systems. D'Limonene reduced the odor from dipentene.

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**EXAMPLE 12**  
**Neutral pH Stripper System #6**

COMPONENT	WEIGHT PERCENT
PROPYLENE CARBONATE	40
N-METHYL PYRROLIDONE	40
D'LIMONENE	5
ISOPROPANOL	5
PD-600	8
METHOCCELL	1
WATER	1

**Procedure:** Same as general method and examples 10 and 11, but methocell and water mixed separately to form microgel before adding to mixture of other components.

**Results:** Composition exhibited good stripping efficiency. Odor masked effectively by the d'limonene.

**EXAMPLE 13**  
**Neutral pH Stripper System #7**

COMPONENT	WEIGHT PERCENT
PROPYLENE CARBONATE	30
N-METHYL PYRROLIDONE	30
DIPENTENE	20
D'LIMONENE	5
ETHANOL	5
PD-600	8
ETHYL CELLULOSE	1
WATER	1

**Procedure:** Same as example 12 and general method.

**Results:** Good stripping efficiency. Ethyl cellulose exhibited better thixotropic characteristics than methyl cellulose.

**EXAMPLE 14**  
**Neutral pH Stripper System #8**

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COMPONENT	WEIGHT PERCENT
PROPYLENE CARBONATE	35
N-METHYL PYRROLIDONE	25
DIPENTENE	20
D'LIMONENE	5
ETHANOL	5
PD-600	8
ETHYL CELLULOSE	1
WATER	1

Procedure: Same as examples 12 and 13 and the general method.  
 Results: Product exhibited good stripping efficiency, no odor.

**EXAMPLE 15**  
Neutral pH Stripper System #9

COMPONENT	WEIGHT PERCENT
PROPYLENE CARBONATE	35
N-METHYL PYRROLIDONE	25
DIPENTENE	15
D'LIMONENE	5
ETHANOL	10
PD-600	8
ETHYL CELLULOSE	1
WATER	1

Procedure: Same as examples 12, 13 and 14 and general method.  
 Results: Product evidenced good stripping efficiency, no odor.

**EXAMPLE 16**  
Neutral pH Stripper System #10

COMPONENT	WEIGHT PERCENT
PROPYLENE CARBONATE	30

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N-METHYL PYRROLIDONE	25
DIPENTENE	15
D'LIMONENE	3
ETHANOL	10
PD-600	5
ETHYL CELLULOSE	1
WATER	1
HCFC-123	10

Procedure: Same as 12, 13, 14 and 15. HCFC-123 added to propylene carbonate, N-methylpyrrolidone, dipentene, d'limonene and surfactant mixture until homogeneous. Thixotropic mixture added to above mixture until homogeneous. Results: Excellent stripping efficiency (about 20-30 minutes). HCFC-123 provided additional stripping efficiency for composition. Odor was masked.

**EXAMPLE 17**  
Neutral pH Stripper System #11

COMPONENT	WEIGHT PERCENT
PROPYLENE CARBONATE	30
N-METHYL PYRROLIDONE	25
DIPENTENE	15
D'LIMONENE	3
ETHANOL	10
ETHYL CELLULOSE	1
PD-600	5
WATER	1
HCFC 113 (FREON)	10

Procedure: Same as Example 16, above.

Results: Excellent stripping efficiency on the order of about 20 to 30 minutes.

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**Example 18**  
**Low pH Stripper Composition #1**

DIPENTENE	40
ISOPROPANOL	25
N-METHYLPYRROLIDONE	25
DODECYLBENZENE SULFONIC ACID (LINEAR)	8
WATER	2

Procedure: Same as general method and examples 1-6.

Results: pH is acidic. Stripping efficiency acceptable (about 60 minutes). Evidences that a composition having a pH below 7 may accelerate stripping action on substrates.

**Example 19**  
**Low pH Stripper Composition #2**

Dipentene	40
Isopropyl alcohol	25
N-Methylpyrrolidone	25
Triethanolamine	4
Dodecylbenzene Sulfonic Acid (linear)	4
Water	2

Procedure: Same as Example 18.

Results: pH is acidic. Stripping efficiency marginal and certainly less acceptable than Example 18. Little stripping action observed for limited timeframe (about 60 minutes). Produced foam when washed off sample plate. Foam was apparently produced by in situ formation of triethanolamine sulfonate.

**Example 20**  
**Low pH Stripper Composition #3**

Dipentene	30
Ethanol	15
N-Methylpyrrolidone	20
Ethylene Carbonate/Propylene Carbonate (50/50)	20
Dodecylbenzene Sulfonic Acid (linear)	8

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## Water

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Procedure: For the most part is the same as Examples 18 and 19. Heated ethylene carbonate sample, added an equal weight of propylene carbonate (solubility of ethylene carbonate is greater than that of propylene carbonate). Ethylene carbonate tended to crystallize out of solution upon cooling- re-solubilized upon heating.

Results: pH is acidic. Flash point lowered by increasing water content. Composition showed stripping activity, but was difficult to work with because of tendency of ethylene carbonate to crystallize out of solution.

Example 21  
Low pH Stripper Composition #4

COMPONENT	WEIGHT PERCENT
Dipentene	20
Ethanol	15
N-Methylpyrrolidone	25
Propylene Carbonate	30
Dodecylbenzene Sulfonic Acid (linear)	4
Triethanolamine	4
Water	2

Procedure: Same as general method and Examples 18 and 19.

Results: Acidic pH. Showed marginal stripping efficiency. Propylene carbonate increased the flash point. Evidence of foam from triethanolamine sulfonate.

Example 22  
Low pH Stripper Composition #5

Component	Weight Percent
Dipentene	20
Ethanol	15
N-Methylpyrrolidone	25
Propylene Carbonate	30

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Dodecylbenzene Sulfonic Acid (linear)	8
Water	2

Procedure: Same as general method and Examples 21, 18 and 19.  
Results: Showed marginal stripping efficiency, perhaps slightly better than example 21, because of absence of triethanolamine sulfonate foam. Propylene carbonate increased the flash point. Removal of triethanolamine resulted in no evidence of foam from triethanolamine sulfonate.

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## EXAMPLE 23

Impregnated Non-Woven Paint Strip Applicator  
Packaged in an Aluminum Foil Barrier Bag

A non-woven, highly absorbent fabric, without chemical binders, thermally bonded composite blend of cellulose and polypropylene- typically a 50-50 (percent by weight) blend is used. Cloth having a basis weight of approximately 2 (oz./yd) has proven satisfactory. Other blends free of materials incompatible with stripper composition may be used. A fabric without additives, fillers or an adhesive promotes stability and shelf life of the product.

The stripper used may be any described hereinabove, but preferred stripper compositions include those as described in Examples 16 and 17.

Barrier bag (one which will not transmit vapor made of aluminum foil), for example, aluminum foil.

Procedure: the fabric bag is slit to an appropriate size, for example 3' by 4', folded and inserted into barrier package sealed on three sides. A metered volume of stripper composition, designed to impregnate the fabric is injected into the open end of the barrier bag which is then immediately sealed.

Results: The applicator is designed to blister and remove coatings from surfaces. It is particularly useful when applied to vertical surfaces because it adheres when applied. The applicator and paint stripper composition is designed so that there is sufficient surface tension between the applicator and the applied surface to retain it and allow it to continuously apply composition, without evaporation, until paint has lifted.

EXAMPLE 24  
Organoclay Composition # 1

Component	Weight Percent	Phase
Bentone EW	2.33	A
H <sub>2</sub> O	13.1	A
Ethanol (95%)	13.1	A

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Dipentene	9.56	B
N-methylpyrrolidone	34.88	B
Propylene Carbonate	24.41	B
PD-600 (Surfactant)	9.56	B

Procedure: Phase A was made first by adding a mixture of ethanol and water to Bentone and then stirring for a period of about 10 to 15 minutes. Note that if water or ethanol is first added and mixed with the Bentone before the other solvent is added, there may be a need to mix the Bentone, H<sub>2</sub>O/ethanol mixture vigorously for a much longer period of time and even then the texture may not be as favorable.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force in a single spindle mixer (about 800 to above 1,000 RPM) for 30 minutes or until homogeneous.

Results: Consistency of the product was similar to petroleum jelly. Stripping efficiency was acceptable (Latex coating removed in about 15 minutes).

EXAMPLE 25  
Organoclay Composition # 2

Component	Weight Percent	Phase
Bentone EW	3.39	A
H <sub>2</sub> O	25.42	A
Ethanol (95%)	8.48	A
N-methylpyrrolidone	33.89	B
Propylene Carbonate	23.73	B
PD-600 (Surfactant)	5.09	B

Procedure: Phase A was made first by adding a mixture of ethanol and water to Bentone and then stirring for a period of about 10 to 15 minutes.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then

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added to phase A under shearing force (about 800 to above 1,000 RPM) for 30 minutes or until homogeneous.

Results: Consistency of the product was similar to petroleum jelly. Stripping efficiency was acceptable.

**EXAMPLE 26**  
Organoclay Composition # 3

Component	Weight Percent	Phase
Bentone EW	7.69	A
H <sub>2</sub> O	9.62	A
Ethanol (95%)	14.42	A
N-methylpyrrolidone	38.46	B
Propylene Carbonate	26.92	B
PD-600 (Surfactant)	2.89	B

Procedure: Phase A was made first by adding a mixture of ethanol and water to Bentone and then stirring for a period of about 10 to 15 minutes.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force (about 800 to above 1,000 RPM) for 30 minutes or until homogeneous.

Results: Consistency of the product was considerably thicker than Examples 23 and 24 and similar to paste. Stripping efficiency was acceptable.

**EXAMPLE 27**  
Organoclay Composition # 4

Component	Weight Percent	Phase
Bentone EW	2.92	A
H <sub>2</sub> O	11.70	A
Ethanol (95%)	11.70	A
Dipentene	11.70	B
N-methylpyrrolidone	29.24	B
Propylene Carbonate	29.24	B

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PD-600 (Surfactant)	3.51	B
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Procedure: Phase A was made first by adding a mixture of ethanol and water to Bentone and then stirring for a period of about 10 to 15 minutes.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force (about 800 to above 1,000 RPM) for 30 minutes or until homogeneous.

Results: Consistency of the product was similar to those of Examples 23 and 24 and to petroleum jelly. Stripping efficiency was acceptable.

**EXAMPLE 28**  
Organoclay Composition # 5

Component	Weight Percent	Phase
Bentone EW	10.31	A
H <sub>2</sub> O	34.35	A
Ethanol (95%)	10.31	A
Dipentene	8.94	B
D'limonene	0.34	B
N-methylpyrrolidone	15.81	B
Propylene Carbonate	13.75	B
PD-600 (Surfactant)	2.75	B
HCFC	3.44	B

Procedure: Phase A was made first by adding a mixture of ethanol and water to Bentone and then stirring for a period of about 10 to 15 minutes. Phase A after mixing was very thick. Amount of Bentone added may necessitate the addition of more ethanol.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force (about 800 to above 1,000 RPM) for about 1 hour or until homogeneous.

Results: Consistency of the product was very thick, similar

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to a very thick paste. Stripping efficiency was less than commercially acceptable, but was evident.

**EXAMPLE 29**  
**Organoclay Composition # 6**

Component	Weight Percent	Phase
Bentone EW	3.16	A
H <sub>2</sub> O	21.05	A
Cabosil™ (Amorphous)	0.52	A
Diatomaceous Earth	0.52	A
Ethanol (95%)	10.53	A
Dipentene	2.11	B
D'limonene	2.11	B
N-methylpyrrolidone	26.32	B
Propylene Carbonate	26.32	B
PD-600 (Surfactant)	3.16	B
HCFC	4.24	B

**Procedure:** Phase A was made first by adding a mixture of ethanol and water to Bentone, Cabosil and diatomaceous earth and then stirring (750 RPM magnetic stirrer) for a period of about 10 to 15 minutes. Phase A after mixing was homogeneous.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force (about 800 to above 1,000 RPM) for about 30 minutes or until homogeneous.

**Results:** Consistency of the product was excellent, quite gel-like and similar to petroleum jelly. Stripping efficiency was quite adequate.

**EXAMPLE 31**  
**Organoclay Composition #7**

Component	Weight Percent	Phase
Bentone EW	4.24	A
H <sub>2</sub> O	24.24	A

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Dipentene	2.43	B
D'Limonene	2.43	B
N-methylpyrrolidone	30.30	B
Propylene Carbonate	30.30	B
PD-600 (Surfactant)	3.64	B
HCFC	2.43	B

Procedure: Phase A was made first by adding water to Bentone, and stirring (750 RPM magnetic stirrer) for a period of about 10 to 15 minutes. Phase A after mixing was homogeneous, but texture was coarse.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force (about 800 to above 1,000 RPM) for about 30 minutes or until homogeneous.

Results: Consistency of the product was good, thick and somewhat gel-like but less gel-like than compositions containing ethanol. Stripping efficiency was quite adequate.

EXAMPLE 32  
Organoclay Composition #8

Component	Weight Percent	Phase
Bentone EW	1.6	A
H <sub>2</sub> O	16.0	A
Ethanol	8.0	A
Dipentene	9.6	B
D'Limonene	1.6	B
N-methylpyrrolidone	32.0	B
Propylene Carbonate	2.4	B
PD-600 (Surfactant)	2.4	B
HCFC	6.4	B

Procedure: Phase A was made first by mixing water and ethanol and then adding the resulting mixture to Bentone. The mixture is stirred (750 RPM magnetic stirrer) for a period of about 10

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to 15 minutes. Phase A after mixing was homogeneous, quite gel-like.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force (about 800 to above 1,000 RPM) for about 30 minutes or until homogeneous.

**Results:** Consistency of the product was good, gel-like, consistency was like petroleum jelly. Stripping efficiency was excellent.

**EXAMPLE 33**  
Organoclay Composition #9

Component	Weight Percent	Phase
Bentone EW	4.0	A
H <sub>2</sub> O	16.0	A
Ethanol	16.0	A
Dipentene	1.6	B
D'Limonene	1.6	B
N-methylpyrrolidone	32.8	B
Propylene Carbonate	22.4	B
PD-600 (Surfactant)	2.4	B
HCFC	3.2	B

**Procedure:** Phase A was made first by mixing water and ethanol and then adding the resulting mixture to Bentone. The mixture is stirred (750 RPM magnetic stirrer) for a period of about 10 to 15 minutes. Phase A after mixing was homogeneous, thick and quite gel-like.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force (about 800 to above 1,000 RPM) for about 30 minutes or until homogeneous.

**Results:** Consistency of the product was good, thick, gel-like, consistency was like a thick petroleum jelly. Stripping efficiency was excellent.

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**EXAMPLE 34**  
Organoclay Composition #10

Component	Weight Percent	Phase
Bentone EW	1.74	A
H <sub>2</sub> O	8.70	A
Ethanol	17.39	A
D'limonene	3.48	B
N-methylpyrrolidone	34.77	B
Propylene Carbonate	24.35	B
PD-600 (Surfactant)	2.61	B
HCFC	6.96	B

**Procedure:** Phase A was made first by mixing water and ethanol and then adding the resulting mixture to Bentone. The mixture is stirred (750 RPM magnetic stirrer) for a period of about 10 to 15 minutes. Phase A after mixing was homogeneous, somewhat thin and quite gel-like.

Phase B was prepared by adding each of the components in any order and then mixing for a period of about 15 minutes in a single spindle mixer at about 800 RPM. Phase B was then added to phase A under shearing force (about 800 to above 1,000 RPM) for about 30 minutes or until homogeneous.

**Results:** Consistency of the product was good, gel-like, consistency was like a thin petroleum jelly. Stripping efficiency was excellent.

**EXAMPLE 35**  
Organoclay Composition #11

Component	Weight Percent	Phase
Bentone EW	3.74	A
H <sub>2</sub> O	27.78	A
Ethanol	6.94	A
D'limonene	1.39	B
N-methylpyrrolidone	18.06	B
Propylene Carbonate	35.42	B

## Claims:

1. A composition for stripping a polymeric coating from a surface comprising:
  - a. about 0% to about 60% by weight of a terpene compound;
  - b. about 10% to about 50% by weight of a compound selected from the group consisting of pyrrolidones and pyrrolidines;
  - c. about 2% to about 20% by weight of a surfactant; and
  - d. about 10% to about 60% by weight of a high flash point, high  $K_B$  solvent selected from the group consisting of ethylene carbonate, propylene carbonate, glycols and glycol ethers.
2. The composition according to claim 1 wherein said solvent is propylene carbonate.
3. The composition according to claim 1 wherein said compound (b) is N-methylpyrrolidone.
4. The composition according to claim 3 further comprising about 5% to about 15% of a solvent extender or bulking agent.
5. The composition according to claim 3 wherein said surfactant is a phosphate ester surfactant.
6. The composition according to claim 5 wherein said surfactant is an alkaline stable mixture of mono and di-substituted phosphate esters of decylalcoholethoxylate.
7. The composition according to claim 5 wherein said surfactant is a phosphate ester of a long-chain alcohol ethoxylate.
8. The composition according to claim 7 wherein said terpene is dipentene.
9. The composition according to claim 9 further comprising an odor-masking compound.
10. The composition according to claim 10 wherein said odor-masking compound is d'limonene.
11. The composition according to claim 5 wherein said solvent extender or bulking agent is ethanol or isopropanol.
12. The composition according to claim 1 additionally comprising between about 0.5% and about 20% by weight water.

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13. The composition according to claim 9 wherein said composition comprises about 1% to about 5% by weight water.

14. The composition according to claim 12 further comprising about 0.02% to about 8% by weight of a thixotropic agent.

15. The composition according to claim 14 wherein said thixotropic agent is selected from the group consisting of cellulose ethers and organoclay rheological agents.

16. The composition according to claim 15 wherein said thixotropic agent is ethyl cellulose and is in the form of a microgel.

17. The composition according to claim 1 further comprising ethanol, water, ethyl cellulose and d'limonene and wherein said terpene is dipentene, said pyrrolidone is N-methylpyrrolidone and said surfactant is a phosphate ester surfactant.

18. The composition according to claim 1 further comprising about 0% to about 8% by weight of an organoclay rheological agent and about 10% to about 45% by weight of a polar solvent selected from the group consisting of water, ethanol, isopropanol and mixtures, thereof.

19. The composition according to claim 18 wherein said organoclay rheological agent is a Bentonet<sup>TM</sup> and said composition is in the form of a gel.

20. The composition according to claim 19 further comprising ethanol and water, said terpene is dipentene, said pyrrolidone is N-methylpyrrolidone and said surfactant is a phosphate ester surfactant.

21. An impregnated applicator for use in stripping a polymeric coating from a surface comprising:

1. a paint stripping composition comprising:

a. about 0% to about 60% by weight of a terpene compound;

b. about 10% to about 50% by weight of a compound selected from the group consisting of pyrrolidones and pyrrolidines;

c. about 2% to about 20% by weight of a surfactant;

d. about 10% to about 60% by weight of a high flash point, high K<sub>B</sub> solvent selected from the group consisting of ethylene carbonate, propylene carbonate, glycols and glycol

ethers; and

2. a fabric piece comprising material stable to said stripping composition, wherein said composition is adsorbed to said fabric piece.

22. The applicator according to claim 21 wherein said fabric piece comprises non-woven, highly absorbent fabric containing an absence of chemical binders.

23. The applicator according to claim 21 wherein said fabric is selected from the group consisting of polyester, cellulose/propylene, rayon/polyester and woodpulp/polyester.

24. The applicator according to claim 21 wherein said solvent (d) is propylene carbonate.

25. The applicator according to claim 21 wherein said compound (b) is N-methylpyrrolidone.

26. The applicator according to claim 21 further comprising about 5% to about 15% of a solvent extender or bulking agent.

27. The applicator according to claim 21 wherein said surfactant is a phosphate ester surfactant.

28. The applicator according to claim 27 wherein said surfactant is an alkaline stable mixture of mono and di-substituted phosphate esters of decylalcoholethoxylate.

29. The applicator according to claim 28 wherein said surfactant is a phosphate ester of a long-chain alcohol ethoxylate.

30. The applicator according to claim 21 wherein said terpene is dipentene.

31. The applicator according to claim 30 further comprising an odor-masking compound.

32. The applicator according to claim 31 wherein said odor-masking compound is d'limonene.

33. The applicator according to claim 26 wherein said solvent extender or bulking agent is ethanol or isopropanol.

34. The applicator according to claim 21 additionally comprising between about 0.5% and about 20% by weight water and about 0.02% to about 8.0% of a thixotropic agent.

35. The applicator according to claim 34 wherein said thixotropic agent is selected from the group consisting of cellulose ethers and organoclay rheological agents.

36. The applicator according to claim 35 wherein said

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thixotropic agent is ethyl cellulose and is in the form of a microgel.

37. The applicator according to claim 21 further comprising about 2% to about 40% by weight of a chlorofluorocarbon.

38. The applicator according to claim 37 wherein said chlorofluorocarbon is selected from the group consisting of  $\text{CHCl}_2\text{CF}_3$  and  $\text{CCl}_2\text{F}_2$ .

39. The applicator according to claim 21 further comprising about 0% to about 8% by weight of an organoclay rheological agent and about 10% to about 45% by weight of a polar solvent selected from the group consisting of water, ethanol, isopropanol and mixtures, thereof.

40. The composition according to claim 37 wherein said organoclay rheological agent is a Bentone<sup>TM</sup> and said composition is a gel.

41. A method of making a paint stripper composition containing a microgel comprising:

1. mixing a thixotropic agent in the presence of water and a solvent extender or bulking agent to produce a microgel-containing suspension; and

2. adding said suspension to a mixture comprising:  
a. about 0% to about 60% by weight of a terpene compound;

b. about 10% to about 50% by weight of a compound selected from the group consisting of pyrrolidones and pyrrolidines;

c. about 2% to about 20% by weight of a terpene emulsifying surfactant; and

d. about 10% to about 60% by weight of a high flash point, high  $K_B$  solvent selected from the group consisting of ethylene carbonate, propylene carbonate, glycols and glycol ethers.

42. The method according to claim 41 wherein said solvent is propylene carbonate.

43. The method according to claim 41 wherein said compound (b) is N-methylpyrrolidone.

44. The method according to claim 41 wherein said surfactant is a phosphate ester surfactant.

45. The method according to claim 44 wherein said

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surfactant is a phosphate ester of a long-chain alcohol ethoxylate.

46. The method according to claim 41 wherein said terpene is dipentene.

47. The method according to claim 41 wherein said solvent extender or bulking agent is ethanol or isopropanol.

48. The method according to claim 41 wherein the amount of water used comprises between about 0.5% and about 20% by weight of the final composition.

49. The method according to claim 41 wherein said thixotropic agent comprises about 0.02% to about 8% by weight of the final composition.

50. The method according to claim 49 wherein said thixotropic agent is selected from the group consisting of cellulose ethers and organoclay rheological additives.

51. The method according to claim 50 wherein said thixotropic agent is ethyl cellulose or a Bentone<sup>TM</sup>.

52. The method according to claim 41 wherein said final composition comprises propylene carbonate, ethanol, water, N-methylpyrrolidone, dipentene, a phosphate ester surfactant and a thixotropic agent.